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64 Contrast agent for NMR Imaging

gay. The agent has improved stability and results in an enhanced water proton relaxation rate. It comprises Sposomes which contain paramagnetic ions bound to physiologically acceptable macromolecules.

SPECIFICATION

	Contrast sgent for NMR imaging	
	The invention relates to novel contrast media for NMR-Medical Imaging. Amongst others the novel contrast media have an improved stability compared with preparations of similar properties; they result in enhanced water proton relaxation rate. The novel contrast media are provided in the form of Eposomes containing peramagnetic ions bound to physiologically acceptable	. ·
10	mecro-molecules. NMR imaging (MRI) is a comparatively new technique which provides a 3-dimensional picture of the human body or of cartain organs thereof in a non-invasive manner. The diagonistic value of TH MRI is greatly enhanced when the proton density information is superimposed on proton relaxation times of tissue water relaxation time information. It is established that the proton relaxation times of tissue water relaxation time information.	10 .
15	reflect not only the composition, and the structural complexity of the decimentation of logical or pathologic state MRI contrast agents are very useful for improving the delineation of structures of organs, for characterizing physiological functions and for the further differentiation	15
20	For this purpose there are generally used paramagnetic lone or stable free reduction. The use of such dramatically shorten water relexation times at relatively-low concentrations. The use of such dramatically shorten water relexation times at relatively-low concentrations. The use of such dramatically shorten water relatively shorten the rest state of the most affective	20
	paramagnetic relexation probes, such as Min' and Go' or stable introduced in the stable of these have not been fully established. The at low dosages. Furthermore the metabolic routes of these have not been fully established. The	25
25	strong complexing agent, such as DTPA, EDITA, BUT this similar large of the Mn ³ -DTPA entrapped in to the blood stream and to blood vessels. Recently the use of the Mn ³ -DTPA entrapped in to the blood stream and to blood vessels. Recently the use of the Mn ³ -DTPA entrapped in to the blood stream and the large through the blood stribution of the metal chelate and	
30	that MMn accumulation did very markedly increase in the speed and in the liver seems reduction in the heart and tudneys relative to free Mn-DTPA. The accumulation in the liver seems to indicate leskage of the complex from the lipersees on their subsequent dissociation.	30
35	of the present invention comprise paramagnetic tons bound to private paramagnetic ions to molecules which are entrapped within liposomes. The binding of the paramagnetic ions to macromolecules enhances the water proton relaxation rate and thus smaller quantities of such ions can be used. This is of importance in view of the substantial toxicity of such ions. The ions can be used. This is of importance in view of the substantial toxicity of such ions. The	35
40	resulting in an extended useful lifetime inside the body. The contrast agents of the invention, resulting in an extended useful lifetime inside the body. The contrast agents of the invention, due to the use of specific liposomes, make possible an improved targeting to specific organs as due to the use of specific liposomes, make possible an improved targeting to specific organs as well as to normal or tumorous tissues. Uposome types developed for targeting drugs to certain organs of the human body can be used for this effect, see for example, Weinstein, UCLA Symp.Mol.Cell Biol. 4, 441 (1983). The peramagnetic ions may be bound to suitable macromosymp.Mol.Cell Biol. 4, 441 (1983). The peramagnetic ions may be bound to suitable macromosymp.	40
45	secules. Mecromolecules of choice are carrain proteins, and secules of the proteins can be used for as to reduce immune reaction problems. The binding properties of the proteins can be used for the bonding of the ions: BSA is known to bind manganese and gandolinium with proton the bonding of the ions: BSA is known to bind manganese and gandolinium with proton the bonding of the ions: BSA is known to bind manganese and gandolinium with proton relaxation enhancement: Blochem 2, 910 (1963) and Blochem 10 (1971), 2834. Experiments relaxation enhancement: Blochem 2, 910 (1963) and Blochem 10 (1971), 2834. Experiments	45
50	well as beta- and gamma-globulins. The experiments have detributed and Mn2* was £8%, solution of such protein dialyzed against 1 mM Mn2*, the fraction of bound Mn2* was £8%, solution of such protein dialyzed against 1 mM Mn2*, the fraction of bound Mn2* was £8%, solution of such proteins, respectively. 53% and 14% respectively for the above defined three types of senum proteins, respectively.	50
	means of a strong complexing agent such as OTPA of EDIA. She has a complexed but the same system can be used with other suitable metal ions. The thus obtained complexes but the same system can be used with other suitable metal ions. The thus obtained complexes but the same system can be used to be a complex inside the	55
85	liposomes does not reduce the relexation effect which seems to be due to the water molecules across the liposome membrane system, thus producing a fast exchange on the NAMR time scale and thus a weighed average of relexation times.	J.
60	described here in detail. See, for example, textbooks such as apparent following Example the vesicles were prepared as set out on Blochemistry 20 833	50
65	(1981). The following Examples are provided in order to illustrate the present invention and they are to be construed in a non-limitative manner. It is clear that a variety of different ions, proteins, cheleting agents and mode of preparation of complexes and liposomes can be resorted to	65
-	The state of the s	

A run was carried out as in Example 1, except that 10% \$-Globulin was used Instead of HSA.

A run was carried out as in Example 1, except that 10% a-Globulin was used instead of HSA.

A run was carried out as in Examples 1-3, but with 1 mM MnCl, instead of 2 mM. Vesicles

25

30

35

Runs were carried out as in Examples 1, 4 and 7, except that HSA-DTPA conjugate replaced

Results of Manganese Binding and Proton Relexation Rates for Liposomes containing Mn? and

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(blank) and in the suspensions of the liposomes, which contained 10% (w/w) of proteins from human sarum. The volume, occupied by the Sposomes, was about 20% of the suspension. The excess manganese concentration in the suspension over thet of the buffer indicates binding of

manganese to the proteins in the vesicles. It is seen from the Table that the largest binding was

85

The measurements were made in two typical frequencies: 21 MHz and 42 MHz, which are

The results of the T, relexation time show a dramatic (up to 33-fold) decreese of T, over that of the blank, which contained menganese in equilibrium with the liposomes. Even when we 60 normalize the results to mangenese concentration, a relexation enhancement of up to factor of . 60

16 is obtained. The best results were obtained for albumin as it binds more Mn2 and it gives

The results for Mn2* and Gd2* bound to protein conjugated with EDTA and OTPA give less 65 relaxation per metal ion, but more metal lons bound per protein. Therefore, the choice between

Corresponding results were obtained with the liposomes containing Gdo.

without departing from the scope and spirit of the invention.

25 containing a corresponding concentration of Mn²⁺ were obtained.

containing this conjugate with the Mn21 were obtained.

Vesicles containing the bound Gd3+ cations were obtained.

containing the conjugate with Mn³⁺ were obtained.

the HSA. Corresponding vesicles were obtained.

45 the HSA. Corresponding vesicles were obtained.

Runs were carried out as in Examples 1 and 4, but with IgG-EDTA conjugate. Vesicles

Runs were carried out as in Examples 1 and 4, but with HSA-EDTA conjugate. Vesicles

A number of runs were carried out as in Examples 1-6, but with Gd Cl₃ replacing MnCl₂.

Runs were carried out as in Examples 1, 4 and 7, except that IgG-DTPA conjugate replaced

In the following there is presented a series of examples of the effects observed: There were measured by atomic absorption manganese ion concentrations in the buffers

washed vesicles, which contain Mn-HSA.

Vesicles were obtained in a similar manner,

Similar vesicles were obtained.

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FYAMPLES EXAMPLE 1:

15 EXAMPLE 2.

EXAMPLE 3:

EXAMPLE 4:

EXAMPLE &

EXAMPLE &

Serum Proteins

55 obtained for the serum albumine.

also large relaxation enhancement.

used in NMR imaging,

35 EXAMPLE 7:

the different systems depends on the perticular application and the clinical results. The T, for suspensions of Sposomes comaining human serum albumin and Mn1* ions at 21 and 42 MHz are given in Table 2. The concentrations of tree manganese ions were kept constant throughout the preparation of the liposomes, including during the process of removal of 5 external proteins. Thus, the additional Mn²° concentrations in the liposome suspensions are due to MnJ* binding to the proteins inside the liposomes. For control experiments we measured T₁ relexation times containing "empty vasicles" i.e. vesicles containing buffer without Mn1*, as well as vesicles containing Mn1* at the same concentration as the outside solutions. Although there was some shortening of T, in these 10 samples compared with the blank solutions, the effect of vasicles containing HSA on T₁ release tion rates is much larger. A comperison to solutions of serum albumin as described in Table 1 should take into consideration the small amount of albumin and bound Mn2* in the suspension of the Eposomes (Table 2). In fact, the normalized effect of the bound Mni*, Tu*'/ \(\Delta \text{Mni*} \) is similar in the two experiments. In an additional experiment which is not described in Table 2 we 15 washed vesicles loaded with 10% HSA and 3mM Mn2* with buffer solution without Mn2*. The results for the total Mn2* concentration in the suspension as measured by stornic absorp tion were [Mn2*]=0.31 mM and T,=48.3 ms at a frequency of 42 MHz. The molar relaxivity, T;"/[Mn11]=69.7 is comparable to the previous experiments. Thus, the fact that the bound manganese was enclosed in Sposomes did not affect its relaxation enhancing properties. It can be concluded that the relaxation obtained in the sytems of the invention is greater by a large factor for the same amount of the toxic, paramagnetic metal ions. Furthermore, toxicity is reduced significantly since the metal ions are entrapped in the Spo-

Contract Contract

somet.

TABLES

		Fraque	Friquency 21 19th		fre	Frequency 42 Mis	1
Samole	150. of		1.8.61	T21, 4 Ha 24	1, .01	1- ا- ۱۰	T10 / 41020
					1		
		3000			3000		•
S S S S S S S S S S S S S S S S S S S	i i	980			1020	•	•
Albusin	•				1180	•	•
a-Globel in	•	20	•		1720	•	•
Y-Glubulla	•	0121	•				•
4010	36.0	156	6.1	9.3	9/1	- S. 4	
	c ;		,	5	6.1	156.	. 66
AI DEBTO	2.34	:	<u>.</u>	: :	×	57.	
4-Globulia	1.62	7.E.		· :	3	11.6	96
8-Globul in	100	1	_				~ *
Alank			1.6	•	Š.	6.9	
1	-	4.2	•	125.	2.0	691	
			123	32.	3.6	S 00	
B. 100019-8	6.5			116	J.	11.2	m
1-610bul 1A	4	d :			25.	18.9	7.2
Slank .	2.64	;			4.3	219.	.99
Albumin	5.93	·		<u>.</u>	;		76
a-Globulin	1.3	4.7	- S				
Y.cloballe	,	. 15.Q	4	105.	19.5	75.	,
			ē	*************			

where $T_{j}(0)$ is the value of T_{j} of an identical solution without a paramagnetic ion.

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,: :•

TABLE 2	Water Praton Spin-Lattice Relaxation Times' in Suspensions of Vesicles	with and without Human Serum Albumin and Mn.
1	Water Praton Spin-Lattice Relat	with and without Huma

Dlack		limpay weiches	Bicks	containing free Mn1.	in in it	Væk	les contain	ing HSA	Vexicles containing 11SA and Mn ^{1*}
[Nn.'.]	7. (sæ)	lata!	F . E	(my).)	T, (ms)	[Ma ¹⁰]	(MA)	T, ⁽ (ms)	T. (13410) (1-1 m.) (1-1)
0.455 0.45 1.86	¥ 5 5	0.545 1,090 2.18*	2 2 4 2 2 4	0.93	120 25 44	0.758 1.213 2.52	0.222 0.195 0.248	* # # =	2 2 8

At NAIR frequency of 42 Mile.

* All edutions contained 130 mar NaCl, 20 mar liepes buffer pal 7.0.

"Venites contained Buffer as in furtnesse b. Mat" was added to the outside solution.

"Verifier prepared by dialisis against solutions identical to those given as Blank.

' Venites prepaind as described in the experimental solution. They were wathed with the solutions given

I F, irlanation tines of the same solutions at a frequency of 21 Mills were 33, 25.5, and 14 ms, respectively. " To he districted between To of the suspensions of verietes with 115A and Ma" and those conterning

hin? eaty. AMn? is the difference of Ma³⁰ concentration in the same two surpression

* Dismeter of weight & standard deviation: 340 ± 74 nm.

. Hismorter of maides t. standard deviations 402 g. 119 nm

10

25

- 1. An MRI contrast enhancer comprising a liposome containing mecromolecule-bound param-CLAIMS egnetic ions.
- 2. An MC1 contrast enhancer according to claim 1 where the paramagnetic ions are selected 5 from Mn3 and Gd3. 3. An MRI contrast enhancer wherein the macromolecules are physiologically acceptable pro-
 - 4. An MRI contrast enhancer according to claims 3, wherein the protein is selected from poins.
- 8. An MRI contrast enhancer according to claim 4, where the serum protein is selected from serum protein.
- serum albumin, beta-globulin and gamma globulin. 6. An MRI contrast enhancer seconding to any of claims 1 to 8, wherein the ions are bound
 - to the protein by absorption forces of the protein. 7. An MRI contrast enhancer according to claims 1 to 5, wherein the paramagnetic lone are
- 15 complexed with a strong complexing agent. 8. An MRI contrast enhancer according to claim 7, where the complexing agent is EDTA or
 - 9. An MRI contrast enhancer according to claims 1 to 8, where the liposome (vesicle) is 8 DTPA.
- phospholipid liposome. 10. An MRI contrast enhancer according to claims 1 to 8, wherein there is used a synthetic
- 11. MRI contrast enhancer systems for use as NMR medical imaging agents, substantially es polymer sposome. hereinbefore described and with reference to any of the Examples.
- 12. An MRI contrast enhancer according to any of claims 1 to 11 in injectable unit dosage form.

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